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MUCIN NUCLEOTIDES

The present invention relates to nucleotide fragments, polypeptides and antibodies and their use in medical treatment and diagnosis.

5 In International Patent Application no. WO-A-88/05054 there is disclosed a tandem repeat sequence contained in the human polymorphic epithelial mucin (HPEM) gene and nucleotide probes, polypeptides, antibodies and antibody-producing cells which are useful in the diagnosis  
10 and treatment of adenocarcinomas such as breast cancer.

The present inventors have now elucidated the nucleotide base sequence of the gene in the region 5' of the tandem repeat sequence (unless the context implies otherwise, directions such as "5'" or "3'", "upstream" or "downstream" used herein refer to the non-template strand of the genomic DNA or fragments thereof). The complete sequence of the 1763 nucleotide bases of the non-template strand upstream of and including the first SmaI restriction site in the tandem repeat is set out in Fig.  
20 1. The sequence of 1575 nucleotide bases of the non-template strand upstream of and including the first SmaI restriction site in the tandem repeat as set out in Fig. 3 has been extended and some parts have been corrected in the light of repeat experiments. The template strand has  
25 a complementary sequence and it is this strand which is transcribed into RNA during expression of the gene

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product.

In addition to conventional transcriptional and translational start sites and intron splicing sites, this sequence contains a number of features which may be  
5 important in the diagnosis and therapy of cancers and in expression of proteins from recombinant vectors. These features will be described below. The amino acid sequence corresponding to the translated portions of this nucleotide sequence gives rise to peptides and thence to  
10 antibodies and antibody-producing cells which may also be useful in such diagnosis and treatment.

In one aspect the present invention provides a nucleic acid fragment comprising a portion of at least 17 contiguous nucleotide bases which portion has a sequence  
15 the same as, or homologous to a portion of corresponding length of the sequence of the coding strand as set out in Fig. 1 or the same as, or homologous to a portion of corresponding length of the sequence complementary to the sequence of the coding strand set out in Fig. 1.

20 As used herein the term "fragment" is intended to include restriction endonuclease-generated nucleic acid molecules and synthetic oligonucleotides.

The nucleic acid fragments of the invention may be single-stranded or double-stranded and they may be RNA or  
25 DNA fragments. Single stranded fragments may be "plus" or coding strands having the sequence of Fig. 1 or a part

thereof or a sequence homologous thereto. Alternatively the single stranded fragments may be "minus" or non-coding strands having a sequence complementary to the sequence of Fig. 1 or a part thereof or a sequence homologous thereto.

5 Double stranded fragments contain a complementary pair of strands, (ie. one plus strand and one minus strand).

RNA fragments according to the invention will, of course, contain uridylic acid ("U") residues in place of the deoxythymidylic acid residues ("T") of the coding 10 (non-template) strand set out in Fig. 1 or, if complementary to the sequence of the coding strand, they will contain U residues in positions complementary to the adenyllic acid ("A") residues in the coding strand set out in Fig. 1.

15 Preferably the nucleic acid fragments of the invention are double-stranded DNA fragments.

Single-stranded nucleic acid fragments of the invention are at least 17 nucleotide bases in length.

Double-stranded nucleic acid fragments of the invention 20 are at least 17 nucleotide base pairs in length.

Preferably the fragments are at least 20 bases or base pairs in length, more preferably at least 25 bases or base pairs and yet more preferably at least 50 bases or base pairs in length.

25 Statistically it is almost certain that a 17 nucleotide base sequence will be unique so that any

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nucleic acid fragment having a contiguous portion of 17 nucleotides of a sequence identical to a portion of corresponding length of the coding strand as set out in Fig. 1, or the same as the non-coding strand complementary 5 to the sequence of Fig. 1, will be new. Fragments according to the invention which are only 17 nucleotides or nucleotide bases in length have a sequence the same as, or complementary to, that set out in Fig. 1. Longer fragments of the invention may have a sequence which is 10 homologous to a corresponding portion of the sequence for the coding strand as set out in Fig. 1 or to the complementary non-coding strand.

Preferably nucleic acid fragments according to the invention have at least 75% sequence homology with a 15 corresponding portion of the sequence of Fig. 1 or the complementary non-coding strand, for instance 80 or 85%, more preferably 90 or even 95% homology. Differences may arise through deletions, insertions or substitutions. In addition to containing a portion homologous to or the 20 same as the sequence of the coding strand in Fig. 1 or complementary non-coding strand, the nucleic acid fragments of the invention may include sequences completely unrelated to that in Fig. 1.

Particular features of interest within the coding 25 strand in Fig. 1 are set out in Tables 1 to 3 below:

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TABLE 1: Signal Sequences

Location*	Sequence in PEM	Significance
5	1-2 CG	transcriptional start site
	73-75 ATG	translational start signal
	131-132 GT	start of first intron
	631-632 AG	end of first intron
10	100-130 } TTCCCTGCTGCTGCT- and } CCTCACAGTGCTTA- 633-637 } CAG...TTGTT	Signal sequence, interrupted by first intron (first intron indicated by "...").
	955-960 CCCGGG	SmaI site at start of tandem repeat

15 Footnotes to Tables 1 and 2

+ In the consensus sequences : R is A or G

N is A, C, G or T

W is A or T

X is

20 Y is C or T

\* Locations are of the 5' base of the indicated PEM  
sequence numbered as in Fig. 1.

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TABLE 2 : Regulatory elements within the 5' flanking sequence

Regulatory element	Consensus Sequence <sup>+</sup>	Sequence in PEM	Location*
SP1	GGGGGG	GGGGGG GGGGGG GGGGGG GGGCGGGCGGGCGGG	-727 -397 -94 -54
SV40 enhancer element			
a	ATGTGTGT	CTGTGGGT	-562
b	GCATGCAT	GCGTGCCT	+25
c	GTGGATAG	GTGGAGAG	-702
AP-1	CTGACTCA G    A	GTGACAC CTGCTTCA GTGCTTAG CTGCTGTA	-739 -418 -61 +27
AP-2	CCCCAGGC G  G	ACCCAGGC CACCGGGC	-597 +77
NF1/CTF	TTGGCTNNNAGCCAA	TTGGCTTTCTCCAA	-618
Glucocorticoid regulatory element:			
Core sequence	TGTTCT	TGTTCT TGTTCC	+38 -321
Consensus sequence	GGTACANNNTGTTCT	GCCIGAAATCTGTTCT AGCIGGCTTTGTTCC	+29 -330
CACCC factor	CACCC	CACCC CACCC	+54 +84
Progesterone receptor consensus sequence	ATTCCCTCIGT	ACTCCTCTCC ACTCCTCTCT ATTTCCTCGGC	-802 -626 -432
Estrogen consensus sequence	GGTCANNNTGACC	GCTCCCGGTGACC	-746
RNA Polymerase III Box A	RRYNNARYXGG	GACCTAGCTGG	-335
Box B	GTTCRANNC	AGTGGAGTGCGG GTTCAGAC	-388 -260
Enhancer sequences:			
Interferon- $\beta$ seq	GGAAATTCTCTCTG	GGAAATTCTCTCC	-642
CMV enhancer	GGAAAGTCCCGTT	GGAAAGTCCGGCT	-585

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The sequence in Fig. 1 also includes two sites occurring in the promoter region and in the first intron having 70 to 80% homology with the mammary consensus, sequence (Rosen, J.M. in "The Mammary Gland, Development, Regulation and Function", Ed. Nevill, M.C. and Daniel, C.W. Plenum Press, pp 301-322). These sites are set out in Table 3 below:

TABLE 3

	Location	Sequence
10		
	-289 to -274	*** * * AGGCTAAACTAGAGC
	+230 to +245	* ** ** GTAAGAATTGCAGACA
15	Consensus	RGAAGRAAANTGGACA

Positions are numbered in accordance with Fig. 1.

\* indicates a mismatch with the consensus sequence.

In the consensus sequence:- R is A or G.

20 N is A, C, G or T.

Preferred fragments according to the present

invention include the transcriptional and translational start signals, "TATAA" box and at least one of the regulatory elements (transcription factor binding sites) set out in Table 2 above. More preferably these fragments contain 2 or 5 more, for instance 3, 4 or 5 of the regulatory elements in addition to the TATAA box or even all of the regulatory elements set out in Table 2. Those fragments containing more than one of the regulatory elements of Table 2 preferably also preserve the relative spacings of those sites from one 10 another and from the TATAA box and transcriptional and translational start signals.

Other preferred fragments of the invention contain at least one of the regions homologous to the mammary consensus sequences as set out in Table 3. Preferably these fragments 15 contain both of the regions having homology with the mammary consensus sequences as set out in Table 3. Those fragments containing both regions having homology with the mammary consensus sequence preferably also preserve the relative spacing of those regions, as found in Fig. 1, from one 20 another and from the TATAA box and transcriptional and translational start signals.

Yet further preferred fragments according to the invention comprise the TATAA box, the transcriptional and translational start signals, at least one and preferably two 25 or more of the regulatory elements as set out in Table 2 and at least one and preferably both of the regions having

homology with the mammary consensus sequence as set out in Table 3. Yet more preferably these fragments also preserve the relative spacing of the features from Tables 1, 2 and 3. Particularly preferred fragments according to the invention

5 comprise the sequence upstream of the TATAA box as set out in Fig. 1 together with, and downstream thereof, transcriptional and translational start signals and a polypeptide coding sequence in correct reading frame register with the promoter sequences and the TATAA box,

10 transcriptional and translational start signals. The coding sequence may encode a part or parts of the polypeptide encoded by the mucin gene, for instance a part or parts thereof other than the tandem repeat sequence, or polypeptides unrelated to that encoded by the mucin gene.

15 Other particularly preferred fragments according to the present invention comprise promoter sequences, a TATAA box, transcriptional and translational start signals and, downstream thereof and in correct reading frame register therewith a coding sequence corresponding to a portion of

20 the mucin gene, for instance corresponding to the first exon (corresponding to bases (1 to 130 of Fig.1.) or a part thereof and/or the second exon (corresponding to bases 633 onwards in Fig.1.) or a part thereof, for instance a part thereof other than the tandem repeat sequence as set out in

25 WO-A-88/05054.

In an especially preferred aspect the fragments

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contain (i) the first 26 bases (bases 1 to 26 of Fig. 1) or (ii) the whole of the first exon (bases 1 to 130 of Fig. 1.) and/or (iii) the splicing/ligating sites for the first intron set out in Table 1 and a non-coding sequence between these 5 sites. The non-coding sequence may be the same as or different to the sequence of the first intron as shown in Fig. 1. Preferably it is the same.

Other preferred fragments of the invention comprise at least a portion of the first intron (bases 231 to 632 of 10 Fig. 1). Further preferred fragments of the invention comprise at least a portion of the 5'-flanking sequence upstream of base -423 of Fig. 1.

Other preferred fragments of the invention comprise a portion of the sequence of Fig. 1 corresponding to a portion 15 of the sequence of Fig. 3.

Further preferred fragments of the invention comprise a combination of any two or more of the foregoing preferred features.

Fragments according to the present invention 20 containing functional coding sequences for a least a part of the first or second exons set out in Fig. 1 are useful in the production of polypeptides corresponding to a part or all of the mucin gene product. Such polypeptides are, in turn useful as immunogenic agents for instance in active 25 immunisation against Human Polymorphic Epithelial Mucin (HPEM) for the prophylactic or therapeutic treatment of

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cancers or raising antibodies for use in passive immunisation and diagnosis of cancers. For use in such methods the fragment, which codes for a polypeptide chain substantially identical to a portion of the mucin core protein, may be 5 extended at either or both the 5' and 3' ends with further coding or non-coding nucleic acid sequence including regulatory and promoter sequences, marker sequences, and splicing or ligating sites. Coding sequences may code for other portions of the mucin core protein chain (for instance, 10 other than the tandem repeat) or for other polypeptide chains. The fragment according to the invention, together with any necessary or desirable flanking sequences is inserted, in an appropriate open reading frame register, into a suitable vector such as a plasmid, or cosmid or a viral 15 genome (for instance vaccinia virus genome) and is then expressed as a polypeptide product by conventional techniques. In one aspect the polypeptide product may be produced by culturing appropriate cells transformed with a vector, harvested and used as an immunogen to induce active 20 immunity against the mucin core protein [Tartaglia et al., Tibtech, 6, 43: (1988)].

Fragments according to the present invention incorporating regulatory elements of Table 2 and/or mammary consensus sequences of Table 3 may be used in securing 25 tissue-specific expression of functional coding sequences in appropriate reading frame register downstream of the

regulatory elements and/or associated with the mammary consensus sequences. Such fragments may therefore be used to express parts or the whole of the mucin gene or any other coding sequence in cells of epithelial origin. Applications 5 of this are in therapy and immunisation where such fragments and associated coding sequences are administered to patients such that the coding sequence will be expressed in epithelial tissues leading to a therapeutic effect or an immune reaction by the patient against the polypeptides.

10 The fragments may be presented as inserts in a vector such as viral genomic nucleic acid and introduced into the patients by inoculation of the vector for instance as a modified virus. The vector then directs expression of the polypeptide *in vivo* and this in turn serves as a therapeutic 15 agent or as an immunogen to induce active immunity against the polypeptide. This strategy may be adopted, for instance, to secure expression of polypeptides encoded by the HPEM gene for treatment or prophylaxis of adenocarcinomas such as breast cancer or to secure tissue 20 specific expression of other peptides under control of the regulatory sequences of Table 1, for instance by administration of a modified vaccinia virus containing the fragment and coding sequences in its genomic DNA. RNA fragments of the invention may similarly be used by 25 administration via a retroviral vector. Selection of tissue specific virus vectors to carry the fragments of the

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invention and coding sequences will further restrict expression of the polypeptide to desired target tissues.

Fragments of the invention may also be used to control expression of oncogenic proteins in experimental 5 transgenic animals. Thus, for instance, a transgenic mouse having an oncogene such as ras, erbB-2 or int 2 expressed under control of the present tissue specific fragments may develop breast tumours and be useful in testing diagnostic agents such as tumour localisation and imaging agents and in 10 testing therapeutic agents such as immunotoxins.

Nucleic acid fragments according to the invention are also useful as hybridisation probes for detecting the presence of DNA or RNA of corresponding sequence in a sample. For use 15 as probes fragments are preferably labelled with a detectable label such as a radionuclide, enzyme label, fluorescent label or other conventional directly or indirectly detectable labels. For some applications, the probes may be bound to a solid support. Labelling of the probes may be achieved by conventional methods such as set 20 out in Matthews *et al.*, *Anal. Biochem.*

169: 1-25 (1988).

In further aspects, the present invention provides cloning vectors and expression vectors containing fragments according to the present invention. The vectors may be, for 25 instance, plasmids, cosmids or viral genomic DNA. The present invention further provides host cells containing

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such cloning and expression vectors, for instance epithelial cells transformed with functional expression vectors containing expressible fragments according to the invention.

The invention further provides nucleic acid fragments, 5 which encode polypeptides as defined below. Such fragments may be fragments as hereinbefore defined. However, in view of the redundancy of the genetic code, nucleic acid sequences which differ slightly or substantially from the sequence of Fig. 2 may nevertheless encode the same 10 polypeptide.

The nucleic acid fragments of the invention may be produced de novo by conventional nucleic acid synthesis techniques or obtained from human epithelial cells by conventional methods, Huynh *et al.*, "DNA Cloning: A 15 Practical Approach" Glover, D.M. (Ed) IRL, Oxford, Vol 1, pp49-78 (1985).

The invention therefore also provides probes, vectors and transformed cells comprising nucleic acid fragments as hereinbefore defined for use in methods of treatment of the 20 human or animal body by surgery or therapy and in diagnostic methods practiced on the human or animal body and for use in the preparation of medicaments for use in such methods. The invention also provides methods for treatment of the human or animal body by surgery or therapy and diagnostic methods 25 practiced in vivo as well as ex vivo and in vitro which comprise administering such fragments, probes, vectors or

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transformed cells in effective non-toxic amount to a human or other mammal in need thereof.

Processes for producing fragments according to the invention and probes, vectors and transformed cells

5 containing them and processes for expressing polypeptides encoded by, or under the regulatory control of, fragments of the invention also form aspects of the invention.

The invention further provides a polypeptide comprising a sequence of at least 5 amino acid residues encoded by the 10 coding portion of the DNA sequence as indicated in Fig. 2. Polypeptides according to the invention preferably have a sequence of at least 10 residues, for instance at least 15, more preferably 20 or more residues and most preferably all the residues shown in Fig. 2.

15 The polypeptide may additionally comprise N-terminal and/or C-terminal sequences not encoded by the DNA sequence indicated by Fig. 2.

20 Polypeptides of the invention containing more than 5 amino acid residues encoded by the DNA sequence in Fig. 2 may include minor variations by way of substitution, deletion or insertion of individual amino acid residues. Preferably such polypeptides differ at not more than 20% 25 preferably not more than 10% and most preferably not more than 5% of residues in a contiguous portion corresponding to a portion of the sequence in Fig. 2.

The invention further provides polypeptides as

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defined above modified by addition of a linkage sugar such as N-acetyl galactosamine on serine and/or threonine residues and polypeptides modified by addition of oligosaccharide moieties to N-acetyl galactosamine or via 5 other linkage sugars. Optionally modified polypeptides linked to carrier proteins such as keyhole limpet haemocyanin, albumen or thyroglobulin are also within the invention.

Polypeptides according to the invention may be 10 produced de novo by synthetic methods or by expression of the appropriate DNA fragments described above by recombinant DNA techniques and expressed without glycosylation in human or non-human cells. Alternatively they may be obtained by deglycosylating native human mucin glycoprotein (which 15 itself may be produced by isolation from samples of human tissue or body fluids or by expression and full processing in a human cell line) [Burchell et al., Cancer Research, 47: 5467-5482, (1987), Gendler et al., P.N.A.S., 84: 6060-6064, (1987)], and digesting the core protein. The polypeptides 20 of the invention are useful in active immunisation of humans, for raising antibodies in animals for use in passive immunisation, diagnostic tests, tumour localisation and, when used in conjunction with a cytotoxic agent, for tumour therapy.

25 The invention further provides antibodies against any of the polypeptides described above.

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As used hereafter the term "antibody" is intended to include polyclonal and monoclonal antibodies and fragments of antibodies bearing antigen binding sites such as the F(ab')<sub>2</sub> fragments as well as such antibodies or fragments thereof which have been modified chemically or genetically in order to vary the amino acid residue sequence of one or more polypeptide chains, to change the species specific and/or isotype specific regions and/or to combine polypeptide chains from different sources. Especially in therapeutic applications it may be appropriate to modify the antibody by coupling the Fab, or complementarity-determining region thereof, to the Fc, or whole framework, region of antibodies derived from the species to be treated (e.g. such that the Fab region of mouse monoclonal antibodies may be administered with a human Fc region to reduce immune response by a human patient) or in order to vary the isotype of the antibody (see EP-A-0 239 400). Such antibodies may be obtained by conventional methods [Williams, Tibtech, 6:36, (1988)] and are useful in diagnostic and therapeutic applications, such as passive immunisation.

The term "antibodies" used herein is further intended to encompass antibody molecules or fragments thereof as defined above produced by recombinant DNA techniques as well as so-called "single domain antibodies" or "dAbs" such as are described by Ward, E.S. et al., Nature, 341:544-546

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(1989) which are produced in recombinant microorganisms, such as Escherichia coli, harboring expressible DNA sequences derived from the DNA encoding the variable domain of an immunoglobulin heavy chain by random mutation 5 introduced, for instance, during polymerise chain reaction amplification of the original DNA. Such dAbs may be produced by screening a library of such randomly mutated DNA sequences and selecting those which enable expression of polypeptides capable of specifically binding the 10 polypeptides of the invention or HPEM core protein.

Antibodies according to the present invention react with HPEM core protein, especially as expressed by colon, lung, ovary and particularly breast carcinomas, but have reduced or no reaction with corresponding fully processed 15 HPEM. In a particular aspect the antibodies react with HPEM core protein but not with fully processed HPEM glycoprotein as produced by the normal lactating human mammary gland.

Antibodies according to the present invention preferably have no significant reaction with the mucin 20 glycoproteins produced by pregnant or lactating mammary epithelial tissues but react with the mucin proteins expressed by mammary epithelial adenocarcinoma cells. These antibodies show a much reduced reaction with benign breast tumours and are therefore useful in diagnosis and 25 localisation of breast cancer as well as in therapeutic methods.

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Further uses of the antibodies include diagnostic tests of assays for detecting and/or assessing the severity of breast, colon, ovary and lung cancers.

The antibodies may be used for other purposes  
5 including screening cell cultures for the polypeptide expression product of the human mammary epithelial mucin gene, or fragments thereof, particularly the nascent expression product. In this case the antibodies may conveniently be polyclonal or monoclonal antibodies.

10 The invention further provides antibodies linked to therapeutically or diagnostically effective ligands. For therapeutic use of the antibodies the ligands are lethal agents to be delivered to cancerous breast or other tissue in order to incapacitate or kill transformed cells. Lethal  
15 agents include toxins, radioisotopes and "direct killing agents" such as components of complement as well as cytotoxic or other drugs.

For diagnostic applications the antibodies may be linked to ligands such as solid supports and detectable  
20 labels such as enzyme labels, chromophores, fluorophores and radioisotopes and other directly or indirectly detectable labels. Preferably monoclonal antibodies are used in diagnosis.

Antibodies according to the present invention may be  
25 produced by inoculation of suitable animals with a polypeptide as hereinbefore described. Monoclonal

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antibodies are produced by known methods, for instance by the method of Kohler & Milstein [Nature, 256: 495-497 (1975)] by immortalising spleen cells from an animal inoculated with the mucin core protein or a fragment 5 thereof, usually by fusion with an immortal cell line (preferably a myeloma cell line), of the same or a different species as the inoculated animal, followed by the appropriate cloning and screening steps.

Antibody-producing cells obtained from animals 10 inoculated with polypeptides of the invention and immortalised such cells form further aspects of the invention.

The invention further provides polypeptides, antibodies and antibody producing cells, such as hybridomas, 15 as hereinbefore defined for use in methods of surgery, therapy or diagnosis practiced on the human or animal body or for use in the production of medicaments for use in such methods. The invention also provides a method of treatment or diagnosis which comprises administering an effective 20 non-toxic amount of a polypeptide or antibody as hereinbefore described to a human or animal in need thereof.

Processes for producing polypeptides according to the invention whether by expression of nucleic acid fragments of the invention or otherwise, and for producing antibodies or 25 fragments thereof and for producing antibody-producing cells such as immortalised cells, form further aspects of the

invention.

The invention further provides a diagnostic test or assay method comprising contacting a sample suspected to contain abnormal human mucin glycoproteins with an antibody 5 as defined above. Such methods include tumour localisation involving administration to the patient of the antibody bearing detectable label or administration of an antibody and, separately, simultaneously or sequentially in either order, administering a labelling entity capable of 10 selectively binding the antibody or fragment thereof. Diagnostic test kits are provided for use in diagnostic tests or assays and comprise antibody and, optionally, suitable labels and other reagents and, especially for use in competitive assays, standard sera.

15 The invention will now be illustrated with reference to the figures of the accompanying drawings in which:

Fig. 1. shows the deoxynucleotide base sequence of the 1763 bases upstream of and including the first SmaI restriction 20 site in the tandem repeat sequence of WO-A-88/05054 using the conventional symbols A, C, G and T for the bases of the non-template strand. The base sequence is arranged in blocks of ten. Untranscribed sequence is in lower case, transcribed sequence is in upper case. The SP1 regulatory 25 elements (Table 2), TATAA box, transcriptional and translational start sites (Table 1) are underlined.

Fig. 2. shows the sequence of the non-template strand commencing from the transcriptional start site, (residue 1 in Fig. 1.) and excluding the sequence of the first intron (bases 131 to 632 of the sequence in Fig.1.). Fig.2 also 5 shows the predicted sequence of the polypeptide using the conventional 1 letter symbols for the amino acid residues. Amino acid residues are numbered down the left-hand side and nucleotide bases down the right hand side. The signal sequence is underlined. The sequences end at the first SmaI 10 site in the tandem repeat.

Fig. 3. shows the deoxy nucleotide base sequence of the 1575 bases upstream of and including the first SmaI restriction site in the tandem repeat sequence of WO-A-88/05054 using 15 the conventional symbols A, C, G and T for the bases of the non-template strand. The base sequence is arranged in blocks of ten in non-coding regions. The exon sequences are shown in blocks of three and translated codons are underlined. The start positions of exons 1 and 2, intron 1 20 and the signal sequence for exon splicing are numbered and labelled. Other features mentioned in Tables 1 and 2 are boxed. The sequence finishes with the first SmaI site of the tandem repeat sequence.

The present invention does not extend to fragments,

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polypeptides and antibodies or related materials such as vectors and cells, which are specifically disclosed in WO-A-88/05054 or WO-A-90/05142, nor to the cDNA fragment whose sequence is indicated in Abe, M. et al., in Biochemical and Biophysical Research Communications, 165(2): 644-649 (1989).

The invention will now be illustrated by the following Examples:

EXAMPLE 1

In an attempt to obtain clones with 5' unique sequences, two gt10 libraries were screened with a probe for the tandem repeat. All the clones obtained lacked any non-repetitive sequence at the 5' terminus. Thus, a different strategy was adopted. To obtain 5' sequence we synthesized the cDNA corresponding to the 5' end of breast cancer cell line transcript using anchored-polymerise chain reaction (A-PCR). The A-PCR procedure [Loh, E.Y. et al., Science, 243: 217-220, (1989)] was used to synthesize cDNA corresponding to the 5' end of the transcript. For the 5' end clones total RNA (5 µg) prepared by the guanidinium isothiocyanate method [Chirgwin, J.M. et al., Biochem., 18: 5294-5299 (1979)] was used for first strand synthesis using a breast cancer cell line (BT20) transcript with AMV-reverse transcriptase (Life Sciences) in a 40 µl reaction mixture

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[Okayama, H. and Berg, P., Mol. Cell. Biol., 2: 161-170 (1982)] containing 1  $\mu$ g of an oligonucleotide primer made to the tandem repeat (5'CCAAGCTTGGAGCCCGGGCCGGCTGGTGTCCGG3'). The total RNA was subjected to reverse transcription, and 5 the products were precipitated with spermine. A poly(dG) tail was introduced with terminal deoxy-transferase (500 U/ml, Pharmacia). Amplification was performed with *Thermus aquaticus* polymerase (Perkin Elmer Cetus) in 100  $\mu$ l of the standard buffer supplied. The primers included the tandem 10 repeat primer and for the poly(dG) end, a mixture of the AN polyC primer (5'GCATGCGCGCGCCGGAGGCCCCCCCCCCCC3') and the AN primer (5'GCATGCGCGCGCCGGAGGCC3') at a ratio of 1:9. Following an initial denaturation at 94°C for 5 min, the reaction was annealed at 55°C for 2 min, extended at 15 72°C for 2.5 min and denatured at 94°C for 1.5 min. Amplification was performed for 30 cycles, and the product was precipitated with ethanol. The DNA was sequentially cut with HindIII and SacII, separated on a 1.2% agarose Gel and the band of approximately 550 bp was purified onto DEAE 20 membrane (Schleicher and Schuell), ligated into pBS-SK<sup>+</sup> and transformed into bacteria XL-1 (Stratagene). This plasmid will be referred to as pBS-5'PEM. All restriction enzymes used were obtained from New England Biolabs Inc., oligonucleotide primers and probes were synthesized on an 25 Applied Biosystems 380B DNA synthesizer.

Four colonies were selected for sequencing, and the

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sequences agreed with each other and with sequence obtained from genomic clones of the region. A Leader sequence of 72 bp preceded the first ATG which was in-frame with the reading frame of the tandem repeat as previously determined 5 (Fig. 1), and the sequence preceding first ATG, CCACCATGA, agrees with the Kozak consensus sequence (Kozak, M., Nucl. Acids. Res., 12: 857-872 (1984)).

The primer extension technique was used to map precisely the position of the capssite. A 21 bp 10 oligonucleotide primer (5'AGACTGGGTGCCCGGTGTCAT3') corresponding to nucleotides 73 to 93 ending at the A of ATG (Fig. 1) was end-labelled with [ $\gamma$ -<sup>32</sup>P]ATP (> 5000 Ci/mmol, Amersham International plc) using T4 poly-nucleotide kinase (Pharmacia) and precipitated three times with equal volumes 15 of 4 M ammonium acetate to remove free [ $\gamma$ -<sup>32</sup>P]ATP from the kinased oligonucleotide. Labelled primer ( $1 \times 10^5$  dpm at  $1 \times 10^7$  dpm/pmol) was annealed to 40  $\mu$ g of total BT 20 RNA in 120 mM sodium chloride at 95°C for 5 min, held at 65°C for 1 h and cooled to room temperature. The annealed primer was 20 extended using 18 units of reverse transcriptase in 50mM Tris pH 8.3 at 45°C, 6 mM magnesium acetate, 10 mM dithiothreitol, 1.8 mM dNTPs in a total volume of 50  $\mu$ l at 45°C for 1h. The reaction was stopped by the addition of 50 mM EDTA and the RNA digested by treatment with RNase-A at 25 400 $\mu$ g/ml for 15 min at 37°C. The samples were then phenol:chloroform extracted prior to ethanol precipitation

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and electrophoresed on a standard 6% sequencing gel yielding two bands which mapped to two C's, 72 and 71 bases upstream of the ATG. The sequencing ladder was single-stranded control DNA (M13mp18) from the Sequenase kit (US Biochemical Corp.).

The most prominent product was 72 bp, equal to the number of base pairs from the 5' end of the oligonucleotide primer to the 5' end of the PCR-derived clone, thus confirming that the cDNA represents the entire length of its corresponding cellular mRNA 5' to the tandem repeat. The presence of a second band may be due to interference with reverse transcriptase by methylation of the C at base 71, since it forms a CpG dinucleotide. Under identical conditions, no primer extension product was seen using RNA from Daudi cells which do not express the PEM mucin.

#### Cloning

A plasmid library, grown in DH1 $\alpha$  cells (RecA-), was used instead of a lambda library, because of the possibility of recombination occurring when lambda is grown in RecA+ cells. This recombination might have been expected, since a part of the tandem repeat sequence (GCTGGGGG) is closely related to the chi sequence (GCTGGTGG) of lambda phage which has been implicated as a hotspot for RecA-mediated recombination in E.coli.

Nucleotide sequence of cDNA clones

Fig 1. shows the DNA sequence from the 5' A-PCR-derived clone, including the consensus sequence of the tandem repeat. Sequences were determined in both 5 directions. The region of conserved tandem repeats was not sequenced in full, although a cDNA tandem repeat clone obtained previously had been circularised, sonicated and about 40 clones sequences [(Gendler *et al.*, *J. Biol. Chem.*, 263:12820-12823 (1988))].

Predicted amino acid sequence and composition of the PEM core protein.

The core protein amino acid composition is dominated  
5 by the amino acid composition of the tandem repeat. Serine,  
threonine, proline, alanine and glycine account for about  
60% of the amino acids.

The deduced sequence of the PEM core protein consists  
of distinct regions including (1) the N-terminal region  
10 containing a hydrophobic signal sequence and degenerate  
tandem repeats and (2) the tandem repeat region itself.  
At the N-terminus a putative signal peptide of 13 amino  
acids follows the first 7 amino acids. However, the actual  
site of cleavage has not been determined as attempts to  
15 obtain N-terminal sequence of the core protein were hindered  
by a blocked amino terminus. Following the signal sequence  
and preceding the first SmaI site (which is used to define  
the beginning of the tandem repeat region) are 107 amino  
acids. Greater than 50% of these amino acids comprise  
20 degenerate tandem repeats. Since the number of tandem  
repeats per molecule is large (greater than 21 for the  
smallest allele we have observed), this domain forms the  
major part of the core protein, and results in a highly  
repetitive structure which is extremely immunogenic.  
25 [Gendler, S. et al., loc. cit]. The sequence of the 20

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amino acid tandem repeat unit corresponds to what might be expected for a protein which is extensively O-glycosylated. Five serines and threonines, four of which are in doublets, are found in the repeat and these potential glycosylation sites are separated by regions rich in prolines (See Fig. 2).

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CLAIMS

1. A nucleic acid fragment comprising a portion of at least 17 contiguous nucleotide bases which portion has a sequence the same as, or homologous to a portion of 5 corresponding length of the sequence of the coding strand as set out in Fig. 1 or the same as, or homologous to a portion of corresponding length of the sequence complementary to the sequence of the coding strand set out in Fig. 1.

2. A fragment according to claim 1 comprising any one or 10 more of the following:

(a) a signal sequence

TTCCTGCTGCTGCTCCTCACAGTGCTTACAGXTTGT

wherein X is an optionally present intron

(b) a mammary consensus sequence AGGCTAAACTAGACCC

15 (c) a mammary consensus sequence GTAAGAATTGCAGACAA

(d) a homologue of a sequence (a), (b) or (c) and

(e) a sequence complementary to a sequence (a), (b), (c) or (d).

3. A hybridisation probe comprising a fragment according 20 to claim 1 or claim 2 bearing a detectable label or linked to a solid support.

4. A cloning or expression vector comprising a fragment

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according to claim 1 or claim 2.

5. A transformed cell comprising a cloning or expression vector according to claim 4.

6. A polypeptide comprising a sequence of at least 5 contiguous acid residues encoded by the coding portion of the DNA sequence as indicated in Fig. 2.

7. An antibody against a polypeptide according to claim 6.

8. An antibody according to claim 7 bearing a detectable label or linked to a solid support.

9. An antibody-producing cell capable of secreting an antibody according to claim 7.

10. A diagnostic kit comprising a fragment according to claim 1 or claim 2 or a probe according to claim 3 or a polypeptide according to claim 6 or an antibody according to claim 7 or claim 8.

11. A fragment according to claim 1 or claim 2 or a probe according to claim 3 or a vector according to claim 4 or a cell according to claim 5 or claim 9 or a polypeptide

according to claim 6 or an antibody according to claim 7 or claim 8 for use in a method of treatment or diagnosis practised on the human or animal body.

12. Use of a fragment according to claim 1 or claim 2 or 5 a probe according to claim 3 or a vector according to claim 4 or a cell according to claim 5 or claim 9 or a polypeptide according to claim 6 or an antibody according to claim 7 or claim 8 in the preparation of a medicament for use in a method of treatment or diagnosis practised on the human or 10 animal body.

13. A method of treatment or diagnosis comprising administering to a cancer patient in need thereof or suspected to have a cancer an effective non-toxic amount of a fragment according to claim 1 or claim 2 or a probe according 15 to claim 3 or a vector according to claim 4 or a cell according to claim 5 or claim 9 or a polypeptide according to claim 6 or an antibody according to claim 7 or claim 8.

14. A method of diagnosis comprising contacting a sample from a patient with a fragment according to claim 1 or claim 20 2 or a probe according to claim 3 or a vector according to claim 4 or a cell according to claim 5 or claim 9 or a polypeptide according to claim 6 or an antibody according to claim 7 or claim 8.

ttactcccttc	cgcccggtcc	gacgcggccc	tcagttcg	cggcccgacc	ccgcggact	cccggtgacc	actagaggc	-753
ggggggggact	ccttggcagt	ggttggaaagt	ggcaaggaaag	gacccttggg	ttcatcgay	cccgaggtttta	ctcccttaag	-703
ttggaaaatttc	ttccccact	cctcttggc	tttcttcaag	gaggggaaacc	aggctgtgg	aaagtccggc	tggggggggg	-603
actgtggtt	caggggaaat	cgggggggg	aacggacag	gggggggtta	gaaaggatggg	gctattccgg	gaagtgtggg	-553
gggggggggg	ccccaaacta	gcacatgtc	cactattat	ccagccctct	tattttctgg	ccgtctctgt	tcgtggacc	-453
gggggggggc	ggggggatgg	ggggggatgg	ctagggtgg	gtttccggac	cttgcgtac	aggacccctcg	ccttagctggc	-403
tttgtttccc	atccccact	tagttgttgc	cctgaggcta	aaactatagc	ccaggggccc	caaggttcccg	actgcccccc	-353
ccccccccc	cgaggccagg	gaglggtgg	tgaagggggg	aggccacgtg	gagaacaaac	gggttagtgcg	ggggtttage	-303
tttttagggcc	tttgttaccc	accaggaaat	ggtttggggag	gaggaggaaag	agttaggagg	taggggggg	gggggggttt	-203
ttgtccatgt	caccgttcg	ctgtgtctcg	ggcgccgggg	ggggggccgt	ataaaagggt	aggccgcgt	-153	
ggcccgctca	CCTCTCAAGC	AGCCAGGCC	TGCCCTGAATC	TGTTCTGCC	CCTCCGCCACC	CATTICACCA	CCACCATGAC	-1+1

Fig. 1

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ACGGGGACCC	CAGTCTCCTT	TCTTCTGCT	GCTGCTCTC	ACAGTCTTAA	CAGgtgggg	gcacgggtg	ggggatggc	107
tgcctgttt	agggtgttt	gtgtgtttt	ctgtggttt	tgctccctgt	cagatggcac	catgaagta	aggtaagaat	+207
tgcagacaga	gctgtccctg	tctgtcccg	aaaggaggaa	aggctaaggaa	caggctgaga	agagtgc	ccaaacctga	+257
gagtgggtac	caggggcaag	caaatagtct	gtagagaatg	ctaggggaa	gagatgtgg	agaggggagg	ctttagaggg	+307
gaaaaaatgc	aggggccatg	agccaaaggcc	tatggcaga	gagaaggagg	ctgtgcagg	gaaaggaggct	tccaaaccag	+357
gggttactga	ggctggccac	tcccgatcc	tccgttattt	attttctctgt	tggccagacg	tttattttc	ttcttgtct	+457
tatttttctt	tcataaagac	ccaaacctat	gacttttaact	tcttacagct	accacagcc	ctaaacccgc	aacagtgttt	+507
ACAGGTCTG	GTCA	TGCAAG	CITACCCCA	GGTGGAGAAA	AGGAGACTTC	GGTACCCAG	AGAAGTTCAG	+557
TACTGAGAAG	AATGCTGTGA	GTATGACCG	CAGGTA	TCCAGCCACA	GGCCGGTTTC	AGGCTCTCC	ACCACTCAGG	+607
GAAGGATGT	CACTTGGCC	CGGGCCAGG	AACCAGCTTC	AGGTTCAGCT	GGCACCTGG	GACAGGATGT	CACCTCGGT	+657
CCAGTCACCA	GGCCAGCCCT	GGGGTCCACC	ACCCGGCAG	CCCAAGATGT	CACCTCAGCC	CGGGACAAACAA	AGCCAGCCCC	+707
GGG								+757

*Fig. 1 Cont'd*

	CCG CTC CAC CTC TCA AGA GCC AGC GCC TGC CTG AAT	36
	CTG TTC TGC CCC CTC GCC ACC CAT TTC ACC ACC ATG ACA CCG GGC ACC OAG TCT CCT	
	T P G T Q S P	96
8	TTC TTC CTG CTG CTG CTC CTC ACA GTG CTT ACA GTT GTT ACA GGT TCT GGT CAT GCA AGC	
	F F L L L L T V L T V V	156
	TCT ACC CCA GGT GGA GAA AAG GAG ACT TCG GCT ACC CAG AGA AGT TCA GTG CCC AGC TCT	
28	S T P G G E K E T S A T Q R S S V P S S	216
	ACT GAG AAG AAT GCT GTG AGT ATG ACC AGC AGC GTACTC RCC AGC OAC AGC CCC GGT TCA	
48	T E K N A V S M T S S V L S S H S P G S	276
	GGC TCC TCC ACC ACT CAG GGA CAG GAT GTC ACT CTG GCC CCG GCC ACG GAA CCA GCT TCA	
68	G S S T T Q G Q D V T L A P A T E P A S	336
	GGT TCA GCT GCC ACC TGG GGA CAG GAT GTC ACC TCG GTC CCA GTC ACC AGG CCA GCC CTG	
88	G S A A T W G Q D V T S V P V T R P A L	396
	GGC TCC ACC ACC CCG CCA GCC CAC GAT GTC ACC TCA GCC CCG GAC AAC AAG CCA GCC CGG	
108	G S T T P P A H D V T S A P D N K P A P	456
	Small GGC	

## SUBSTITUTE SHEET

Fig. 2

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'608 TTGCTCTCC AAGAAGGAA CCCAGGTCCG TGAAAGTCCG GCTGGGCGG ACTGTTGGTT TACGGGTAGA  
 537 AGGGCTGTG GAACGAAACG GGAGGGTTA GAAGGGTGGG GCTATTCGGG AAGTGTGG GGGAGGGAGC  
 '467 CCAAAACTAG CACCTAGTCC ACTCATTAATC AAACCGCTT ATTCTGGC CGCTCTGCTT CAGTGGACCC  
 '388 GGGAGGGGC GGGAAAGTGG AGTGGGAGAC CTAGGGTGG GCTTCCGAC CTTCGCTAC AGGACCTCGA  
 '327 OCTTAGCTGGC TTGTGTCGC ATCCGCACGT TAGTTGTCG CCTGAGGCTA AACTAGCC GAAGGGCC  
 '257 CAAAGTCCAG ACTGCGCTCC CCCTCCCGG AGCCAGGGAG TGGTGGTGA AAGGGAGGC CAGCTGGAGA  
 '187 ACAAAAGGGT AGTCAGGGGG TTGAGGAGTT AGAGGCCCTG TACCTACCC AGGAATGGTT GGGAGGGGA  
 '117 GGAAGAGGTA GGAGGTAGGG GAGGGGGCGG GGTTCGGTCA CCTGTCACCT GCTCGCTGTG CCTAGGGCGG  
 '467

Fig.3

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47 GGGGGGGG AGTGGGGGA CGGTATAAA GCGGTAGGGC CCTGTGC 1 CCG CTC CAC CTC TCA ACA GCC  
 AGG GGC TGC CTC AAT CTG TTC TGC CCC CTC CCC ACC CAT TTC ACC ACC ATG ACA CCG GGC ACC CAG 73  
 97 SIGNAL SEQUENCE  
 TCT CCTCTCG CTG CTCA GTG CTA GAG 131 INTRON 1  
 GAGGTGGGG AGTTGGGCTT GCCCTTGCTT AGGTGGCTT TGGTGGTTC TTTCTGTGGG CTTTTGCTCC  
 CTGGAGATG GCACCATGAA GTTAAGGTA GAATATCAGA CAGGGCTGC CCTGCTGTG CCAGAAGGAG  
 GGAGGGCTA AGGACAGGCT GAGAAAGATT GCCCCCCAACCT CTGAGAGTGG GTACCCAGGGG CAAGCAAATG  
 TCCTGTAGAG AAGTCAGGG GGAAGAGT AGGGAGGG AAGGCTTAAG AGGGAAAGAA ATGCAAGGGC  
 CATGAGCCAA GGGCTATGGC CAGAGGAG GAGGCTGCTG CAGGGAAAGGA GGC TTCAAAC CCAGGGTTA

Fig. 3 *Cont'd(1)*

SUBSTITUTE SHEET

CTGAGGCTGC CCACCTCCCA GTCCCTCTGG TATTATTCTT CTGGTGGCCA GAGCTTATAT TTTCTTCTTG  
 CTCTTATTTT TCCCTCAAA AGACCCAACC CTATGACTTT AACTCTTAC AGCTTACACA GCCCTAAAC

640 641 EXON 2  
 CCGCAACAG TT GTT ACA GGT TCI GGT CAI GCA AGC TCI ACC CCA GGT GGA GAA AAG GAG  
 ACT TGG GCT ACC AGG AGA AGT TCA GTC CCC AGG TCI ACT GAG AAT GCT GIG AGT ATG ACC AGC AGC

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 GTA CTC AGC CAC AGC CCC GGT TCA GGC TCC CCA CCA CTC AGG GAC AGG AIG TCA CTC TGG CCC CGG  
 CCA CGG AAC CAG CTT CAG GTC CCA CGT GGT GGG GAC AGG AIG TCA CTT CGG TCC CAG TCA CCA CGA

968  
 GCC CTG GGC TCC ACC ACC GGG CCA GCC CAC GAT GTC ACC TCA GCC CCG GAC AAC AAG CCA GCG CGG

→ TANDEM REPEAT

Fig. 3 Cont'd(2)

## INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 90/02020

## I. CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all) \*

According to International Patent Classification (IPC) or to both National Classification and IPC

IPC<sup>5</sup>: C 07 H 21/04, C 07 K 13/00, C 12 N 15/12, C 12 P 21/00,  
C 12 Q 1/68, G 01 N 33/574, C 07 K 15/00

## II. FIELDS SEARCHED

Minimum Documentation Searched ?

Classification System	Classification Symbols
IPC <sup>5</sup>	C 07 K, C 12 N, C 12 P, C 12 Q
Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched *	

## III. DOCUMENTS CONSIDERED TO BE RELEVANT \*

Category *	Citation of Document, <sup>11</sup> with indication, where appropriate, of the relevant passages <sup>12</sup>	Relevant to Claim No. <sup>13</sup>
X	WO, A, 8805054 (IMPERIAL CANCER RESEARCH TECHNOLOGY) 14 July 1988 see the whole document; especially claims cited in the application	6
Y	--	3-5,7-10,14
P,X	Journal of Biological Chemistry, vol. 265, no. 10, 5 April 1990, The American Society for Biochemistry and Molecular Biology, Inc., (US), M.J.L. Ligtenberg et al.: "Episialin, a carcinoma-associated mucin, is generated by a polymorphic gene encoding splice variants with alternative amino termini", pages 5573- 5578 see the whole article	1,2,6-9
P,Y	--	3-5,10,14

\* Special categories of cited documents: <sup>10</sup>"A" document defining the general state of the art which is not  
considered to be of particular relevance"E" earlier document but published on or after the International  
filing date"L" document which may throw doubt on priority claim(s) or  
which is cited to establish the priority date of another  
document which is cited for its special relevance (as specified)"O" document referring to an oral disclosure, use, exhibition or  
other means"P" document published prior to the International filing date but  
later than the priority date claimed"T" later document published after the International filing date  
or priority date and not in conflict with the application but  
used to understand the principle or theory underlying the  
invention"X" document of particular relevance; the claimed invention  
cannot be considered novel or cannot be considered to  
involve an inventive step"Y" document of particular relevance; the claimed invention  
cannot be considered to involve an inventive step when the  
document is combined with one or more other such docu-  
ments, such combination being obvious to a person skilled  
in the art

"a" document member of the same patent family

## IV. CERTIFICATION

Date of the Actual Completion of the International Search

3rd April 1991

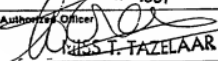
Date of Mailing of this International Search Report

16 MAY 1991

International Searching Authority

EUROPEAN PATENT OFFICE

Signature of Authorized Officer


  
HANS T. TAZELAAR

III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)		
Category *	Citation of Document, <sup>11</sup> with indication, where appropriate, of the relevant passage	Relevant to Claim No.
P,X	Journal of Biological Chemistry, vol. 265, no. 25, 5 September 1990, The American Society for Biochemistry and Molecular Biology, Inc., (US), S.J. Gendler et al.: "Molecular cloning and expression of human tumor-associated polymorphic epithelial mucin", pages 15286-15293 see figure 1 --	1,2,6
P,X	Journal of Biological Chemistry, vol. 265, no. 25, 5 September 1990, The American Society for Biochemistry and Molecular Biology, Inc., (US), M.S. Lan et al.: "Cloning and sequencing of a human pancreatic tumor mucin cDNA", pages 15294-15299 see the whole article --	1,2,6-9
P,A	WO, A, 9005142 (IMPERIAL CANCER RESEARCH TECHNOLOGY) 17 May 1990 see abstract and claims cited in the application -----	1-10,14

## FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET

V.  OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE<sup>1</sup>

This International search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons:

1.  Claim numbers XX....., because they relate to subject matter not required to be searched by this Authority, namely:  
Claims 11-13

Pls. see Rule 39.1 (iv) - PCT:

Methods for treatment of the human or animal body by surgery or therapy, as well as diagnostic methods.

2.  Claim numbers....., because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3.  Claim numbers....., because they are dependent claims and are not drafted in accordance with the second and third sentences of PCT Rule 6.4(a).

VI.  OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING<sup>2</sup>

This International Searching Authority found multiple inventions in this International application as follows:

1.  As all required additional search fees were timely paid by the applicant, this International search report covers all searchable claims of the International application.

2.  As only some of the required additional search fees were timely paid by the applicant, this International search report covers only those claims of the International application for which fees were paid, specifically claims:

3.  No required additional search fees were timely paid by the applicant. Consequently, this International search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:

4.  As all searchable claims could be searched without effort justifying an additional fee, the International Searching Authority did not invite payment of any additional fee.

## Remark on Protest

The additional search fees were accompanied by applicant's protest.  
 No protest accompanied the payment of additional search fees.

ANNEX TO THE INTERNATIONAL SEARCH REPORT  
ON INTERNATIONAL PATENT APPLICATION NO.

GB 9002020  
SA 43255

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 07/05/91. The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Patent family member(s)			Publication date
WO-A- 8805054	14-07-88	AU-A-	1103988	27-07-88	
		EP-A-	0341252	15-11-89	
		JP-T-	2501828	21-06-90	
WO-A- 9005142	17-05-90	None			